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Genome Evolution in the Genus *Sorghum* (Poaceae)

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• **Background and Aims** The roles of variation in DNA content in plant evolution and adaptation remain a major biological enigma. Chromosome number and 2C DNA content were determined for 21 of the 25 species of the genus *Sorghum* and analysed from a phylogenetic perspective.

• **Methods** DNA content was determined by flow cytometry. A *Sorghum* phylogeny was constructed based on combined nuclear ITS and chloroplast *ndhF* DNA sequences.

• **Key Results** Chromosome counts ($2n = 10, 20, 30, 40$) were, with few exceptions, concordant with published numbers. New chromosome numbers were obtained for *S. amplexum* ($2n = 30$) and *S. leiocladum* ($2n = 10$). 2C DNA content varies 8.1-fold (1.27–10.30 pg) among the 21 *Sorghum* species. 2C DNA content varies 3.6-fold from 1.27 pg to 4.60 pg among the $2n = 10$ species and 5.8-fold (1.52–8.79 pg) among the $2n = 20$ species. The $x = 5$ genome size varies over an 8.8-fold range from 0.26 pg to 2.30 pg. The mean 2C DNA content of perennial species (6.20 pg) is significantly greater than the mean (2.92 pg) of the annuals. Among the 21 species studied, the mean $x = 5$ genome size of annuals (1.15 pg) and of perennials (1.29 pg) is not significantly different. Statistical analysis of Australian species showed: (a) mean 2C DNA content of annual (2.89 pg) and perennial (7.73 pg) species is significantly different; (b) mean $x = 5$ genome size of perennials (1.66 pg) is significantly greater than that of the annuals (1.09 pg); (c) the mean maximum latitude at which perennial species grow (–25.4 degrees) is significantly greater than the mean maximum latitude (–17.6) at which annual species grow.

• **Conclusions** The DNA sequence phylogeny splits *Sorghum* into two lineages, one comprising the $2n = 10$ species with large genomes and their polyploid relatives, and the other with the $2n = 20, 40$ species with relatively small genomes. An apparent phylogenetic reduction in genome size has occurred in the $2n = 10$ lineage. Genome size evolution in the genus *Sorghum* apparently did not involve a ‘one way ticket to genomic obesity’ as has been proposed for the grasses.

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Key words: *Sorghum bicolor*, wild sorghum, genome size, DNA content, chromosome numbers, systematics.

INTRODUCTION

Sorghum bicolor (L.) Moench, sorghum, is one of the world's major grain crops. It is extensively cultivated in marginal rainfall areas of the tropics and subtropics, and selected varieties are widely grown in temperate climates. The wild species of sorghum represent a potential diverse source of germplasm for sorghum breeding programmes. The genus *Sorghum* has 25 recognized species that have been taxonomically classified into five subgenera or sections: *Eusorghum*, *Chaetosorghum*, *Heterosorghum*, *Parasorghum* and *Stiposorghum* (Garber, 1950). Species of the genus have chromosome numbers of $2n = 10, 20, 30$ or 40 (Garber, 1950; Lazarides *et al.*, 1991).

Section *Eusorghum* includes cultivated sorghum *S. bicolor* and its subspecies *drummondii* and *arundinaceum*, and the wild species *S. alnum* Parodi, *S. propinquum* (Kunth) Hitch. and *S. halepense* (L.) Pers. (deWet, 1978). Species of section *Eusorghum* have a natural range through Africa and southern Asia (deWet, 1978; Duvall and Doebley, 1990). *Sorghum bicolor* and *S. propinquum* are $2n = 20$ species that are chromosomally similar. Hybrids of *S. bicolor* and *S. propinquum* are meiotically regular with

ten bivalents observed at meiotic metaphase I (Doggett, 1988). *Sorghum halepense* ($2n = 40$) is polyploid. Meiotic analysis of *S. bicolor* \times *S. halepense* hybrids revealed *S. halepense* to possess one genome very similar to *S. bicolor*, and another more divergent or rearranged genome, thus suggesting that *S. halepense* is a disomic polyploid (allopolyploid) or perhaps a ‘segmental allopolyploid’ (Duara and Stebbins, 1952; Endrizzi, 1957; Tang and Liang, 1988).

Section *Parasorghum* comprises seven Asian, Australian and central American species. Sections *Chaetosorghum* and *Heterosorghum* are monotypic and native to the Australo-Pacific region. The ten species of section *Stiposorghum* are found in northern Australia (Lazarides *et al.*, 1991).

Phylogenies based on sequence analysis suggest that the *Sorghum* section designations may not correspond to evolutionary relationships (Spangler *et al.*, 1999; Dillon *et al.*, 2001). Recent sequence and systematic data have led Spangler (2003) to split *Sorghum* into three genera, *Sorghum*, *Sarga* and *Vacoparis*. However, the limitations of the available sequence-based phylogenies suggest that this reclassification is premature.

As part of a larger analysis of *Sorghum* systematics, phylogeny, genomics and germplasm, the DNA content

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has been determined and chromosome numbers obtained of 21 *Sorghum* species. The data acquired are discussed in an evolutionary and phylogenetic context.

METHODS

Plant material

The name, herbarium voucher number, accession number, life form and origin of the *Sorghum* species analysed for DNA content and chromosome number are listed in Table 1. All plants were grown from seeds in a glasshouse, except for *S. halepense* that was sampled directly from the field. The geographic and latitudinal distributions of native Australian *Sorghum* species were taken from Lazarides *et al.* (1991) and herbarium records of the Australian Tropical Crops and Forages Collection, Queensland.

Chromosomes

Chromosome counts were made with slight modification by the protocol of Jewell and Islam-Faridi (1994). Root-tips (approx. 4 mm long) were removed from plants and treated with an aqueous 0.4 % 8-hydroxyquinoline solution for 5 h at room temperature, fixed in 95 % ethanol–glacial acetic acid (4 : 1 v/v), rinsed several times with distilled water, hydrolysed for 5 min in 0.1 N HCl, rinsed 5 min with distilled water, and washed in citrate buffer (pH 4.5) for 5 min. To digest the cell wall, root-tips were treated for 15–50 min at 37 °C with pH 4.5 aqueous 5 % cellulase (Onozuka R-10; Yakult Honsha Co. Ltd, Tokyo) and 1.0 % pectolyase Y-23 (Seishin Corporation, Tokyo), and rinsed three times with distilled water. Rinsed meristems were placed on a clean glass slide with a drop of ethanol–acetic acid (3 : 1), teased apart with a fine-tipped pair of tweezers, and allowed to air dry at room temperature for 2 d. The chromosomes were stained with Azure Blue. Chromosomes from two or more plants of each accession were counted.

Determination of DNA content

At least three plants for each species were analysed to obtain the mean DNA content. Newly expanded leaves of the target species and of a standard species were manually diced together in buffer (Galbraith *et al.*, 1983), pH 7.2, to release nuclei as described by Johnston *et al.* (1999). The diced leaves were filtered through a 53- μ m nylon mesh, and propidium iodide was added to a final concentration of 50 p.p.m. The mean fluorescence of nuclei was quantified using a Coulter Epics Elite flow cytometer (Coulter Electronics, Hialeah, FL) equipped with a water-cooled laser tuned at 514 nm and 500 mW. Fluorescence at >615 nm was detected with a photomultiplier screened by a long pass filter. Mean 2C DNA content of each target species was calculated by comparing its mean nuclear fluorescence with the mean fluorescence of the nuclei of an internal standard. Due to the range and distribution of *Sorghum* DNA contents, one of three different internal standards was used to avoid overlap of the standard and target species. One

standard, *Arabidopsis thaliana* ecotype Columbia has a genome size of 157 Mb or 1C = 0.16 pg (Bennett *et al.*, 2003). The DNA content of the other standards, *S. bicolor* Tx623 (2C DNA content = 1.67 pg) and *Lupinus texensis* (2C DNA content = 2.44 pg), was determined from 15 replicates of diced mixtures of leaves from *S. bicolor*, *L. texensis* and *A. thaliana* Columbia, where *A. thaliana* (2C = 0.32 pg, 1C = 157 Mbp; Bennett *et al.*, 2003) was the calibration standard.

Sequence phylogeny

A phylogenetic tree based on combined ITS/*ndhF* DNA sequences was constructed to evaluate the evolution of chromosome number and DNA content. The DNA sequences used were from Dillon *et al.* (2001) and S. L. Dillon (pers. com.) with all sequences available from Genbank. Genbank accession numbers for ITS1 are: AF302909–AF302913; AF302915–AF302918; AF302920–AF302921; AF302924–AF302927; AY048867; AY048871; AY282488; AY282490; UO4793; UO4795; U46612. Genbank accession numbers for *ndhF* are: AF117423–AF117424; AF117426; AF117430–AF117432; AY048873–AY048874; AY282470–AY282475; AY282477–AY282481; AY282484; U21981; U21985.

Forward and reverse sequences were assembled and edited using Sequencher 3.0 (Gene Codes Corp.). The lengths of the aligned *ndhF* and ITS1 sequences were 2014 and 252 nucleotides, respectively. Sequences were aligned utilizing CLUSTAL X (improved version of CLUSTAL V; Higgins *et al.*, 1992) using a gap opening of 10, a gap extension of 0.5 and a transition weight of 0.5. *Zea mays* sequences were included as an outgroup species. Phylogenetic analyses were carried out on the *ndhF*/ITS1 dataset using PAUP* 4.0b2 (Swofford, 1999). Prior to combining the ITS1 and *ndhF* data sets, a congruence test (partition-homogeneity test in PAUP) was used to determine if there was significant conflict between the data sets. This test compared the sum of lengths of the most parsimonious trees to the distribution of the sum of lengths of the most parsimonious trees using random partition of the characters, 100 reps, and tree bisection–reconnection. Since there were no significant amounts of conflict ($P < 0.01$) between the ITS1 and *ndhF* data sets, they were combined for analysis. A maximum parsimony branch and bound search was performed with all characters having equal weight and gaps treated as missing. Branches with minimum length of zero were collapsed, and duplicate trees were eliminated from the set of most parsimonious trees. There were 66 phylogenetically informative characters. Bootstrap data were generated using the fast stepwise addition for 10 000 replicates with TBR branch swapping and multitrees option in effect.

RESULTS

Chromosomes

Chromosome numbers of the 21 sorghum species studied are listed in Table 2. Figure 1 shows karyotypes of *Sorghum* species varying in chromosome number and

TABLE 1. Accession number, life form and origin of 21 *Sorghum* species analysed for DNA content and chromosome number

Species	Herbarium voucher	Accession no. ¹	Life form	Collection date and site, or source of seeds
<i>Sorghum amplum</i> Lazarides	CANB 480260 ²	302455 ²	Annual	17 Mar. 1994, 1.4 km E of Lake Argyle/T/O on Great Northern Hwy, WA, Australia
<i>S. angustum</i> S. T. Blake	BRI AQ585981 ³	302605 ³	Annual	19 May 1995, Windmill Ck crossing, 18.8 km S of Musgrave, Station on Peninsula. Development Road, QLD, Australia
<i>S. bicolor</i> TX623 L. (Moench)			Annual	Seeds obtained from W. Rooney, Texas A&M University
<i>S. bicolor</i> Pioneer 8695 L. (Moench)			Annual	Seeds obtained from L. Rayburn, University of Illinois
<i>S. brachypodium</i> Lazarides	CANB 480293	302481	Annual	19 Mar. 1994, Jabiru Ring Road East, Jabiru, NT, Australia
<i>S. bulbosum</i> Lazarides	DNA D129483 ⁴	302645	Annual	25 Apr. 1996, 29.1 km S Wyndham T/O on Halls Creek Rd (Great Northern Hwy) NT, Australia
<i>S. ecarinatum</i> Lazarides	DNA D129486	302661	Annual	1 May 1996, 10.1 km NW of Windjana T/O on Gibb River Rd, Napier Range, Napier Creek, NT, Australia
<i>S. exstans</i> Lazarides	BRI AQ586005	302577	Annual	2 Apr. 1995, 37 km N Pickataramoor on Melville Island, Australia
<i>S. halepense</i> (L.) Pers.			Perennial	14 June 2001, Hwy 60, 0.5 miles W of Brazos River, TAES Field Laboratory, Burleson County, TX, USA
<i>S. interjectum</i> Lazarides	JC 2087 ⁵	302445	Perennial	54.1 km W of Halls Ck on Great Northern Hwy, WA, Australia
<i>S. intrans</i> F. Muell. Ex Benth.	JC 2023	302389	Annual	11 Mar. 1994, Hotel at Haynes Creek on Stuart Hwy, NT, Australia
<i>S. laxiflorum</i> Bailey	BRI AQ773635	302510	Annual	15 Apr. 1994, 67.9 km N of Wollogorang on Wollogorang Station Rd to coast, NT, Australia
<i>S. leiocladum</i> (Hack.) C E. Hubb	DNA D0155521	300170	Perennial	16 Dec. 1997, 2–3 km W from Drake on roadside on range in State Forest, NSW, Australia
<i>S. macrospermum</i> Garber	DNA C867	302367	Annual	4 Apr. 1995, 7.9 km N Katherine River bridge on Stuart Hwy, NT, Australia
<i>S. matarankense</i> Garber & Snyder	DNA D129480	302637	Annual	22 Apr. 1996, 13.8 km W Carpenteria Hwy T/O on Stuart Hwy, NT, Australia
<i>S. nitidum</i> (Vahl.) Pers.	CANB 479893	302539	Perennial	21 Apr. 1994, Paddock T330 CSIRO Lansdown Res. Stn on Flinders Hwy, QLD, Australia
	JC 2218	302540	Perennial	21 Apr. 1994, 22.7 km SE of Woodstock on Giru Woodstock Rd, QLD, Australia
	BRI AQ740677	302542	Perennial	21 Apr. 1994, 10.5 km E of Ayr on Alva Beach Rd, QLD, Australia
	CANB 479881	302543	Perennial	22 Apr. 1994, 0.5 km S of Mt Stuart lookout on Mt Stuart Rd, Townsville, QLD, Australia
	ATCGRC 0007 ⁶	302558	Perennial	18 May 1994, 6.9 km N of Byfield on Byfield Rd, QLD, Australia
	BRI AQ773677	302559	Perennial	23 May 1994, 1.6 km N of Bakers Rd, 14 km N of Yeppoon T/O on Bruce Hwy, QLD, Australia
	BRI AQ496360	316930	Perennial	6 Jun. 2000, 450 m down road from summit of Mt Stuart on both sides of road, QLD, Australia
<i>S. plumosum</i> (R. Br.) P. Beauv.	BRI AQ773634	302489	Perennial	11 Apr. 1994, Einslie River, 26.4 km W of Georgetown on Gulf Development Rd, QLD, Australia
	DNA D129468	302635	Perennial	21.5 km N 3-Ways T/O on Stuart Hwy, NT, Australia
<i>S. propinquum</i> (Kunth) Hitch.			Perennial	Africa, seeds from W. Rooney, Texas A&M University
<i>S. purpureosericeum</i> (A. Rich). Aschers & Schweinf	IS 18945 ⁷	318068	Annual	Sudan, NE tropical Africa
<i>S. stipoides</i> (Ewart & Jean White) C. Gardner and C. E. Hubb	DNA D129465	302614	Annual	2 May 1996 102 km N of Barnett River Roadhouse on Gibb River Rd, NT, Australia
<i>S. timorense</i> (Kunth) Buse	DNA D129474	302660	Annual	1 May 1996, 5.7 km NW of Windjana T/O on Gibb River Rd, NT, Australia
<i>S. versicolor</i> (2x) Anderss.			Annual	East Africa, seeds obtained from G. Liang, Kansas State University
<i>S. versicolor</i> (4x) Anderss.			Annual	Colchicine-induced autotetraploid by G. Liang, Kansas State University

¹ AusTRC number, Australian Tropical Crops and Forages Collection, Queensland Department of Primary Industries;² CANB, Australian National Herbarium, Canberra, ACT Australia;³ BRI, Queensland Herbarium, Mt Coot-tha, QLD, Australia;⁴ Northern Territory Herbarium, Darwin, NT, Australia;⁵ JC, Jeff Corfield Collection, Townsville, QLD, Australia;⁶ ATCGRC, Australian Tropical Crops Genetic Resource Centre Collection;⁷ IS, ICRISAT.

TABLE 2. DNA content and chromosome number of 21 *Sorghum* species

Species	2C DNA content			Duncan grouping ²	$x = 5$ genome size (pg) ³	Reported chromosome no. ⁴ ($2n$)	Counted chromosome no. ($2n$)	Section
	pg	SE	Mbp/1C ¹					
<i>Sorghum amplus</i> ⁵	7.69	0.032	3768	C	1.28	10	30	Stiposorghum
<i>S. angustum</i> ⁵	3.70	0.042	1813	H	1.85	10	10	Stiposorghum
<i>S. bicolor</i> ⁵ (Tx623)	1.67	0.007	818	M	0.42	20	20	Eusorghum
<i>S. bicolor</i> ⁵ (Pioneer 8695)	1.67	0.012	818	M	0.42	20	20	Eusorghum
<i>S. brachypodium</i> ⁵	3.36	0.129	1646	I	1.68	10	10	Stiposorghum
<i>S. bulbosum</i> ⁶	2.30	0.009	1127	K L	1.15	10	10	Stiposorghum
<i>S. ecarinatum</i> ⁶	2.10	0.002	1029	L	1.05	10	10	Stiposorghum
<i>S. exstans</i> ⁵	2.75	0.006	1348	J	1.38	10	10	Stiposorghum
<i>S. halepense</i> ⁵	3.28	0.009	1607	I	0.41	40	40	Eusorghum
<i>S. halepense</i> ⁷	3.27	0.020	1602	I	0.41	40	40	Eusorghum
<i>S. interjectum</i> ⁷	7.29	0.048	3572	D	1.22	30	30	Stiposorghum
<i>S. intrans</i> ⁶	2.28	0.023	1117	K L	1.14	10	10	Stiposorghum
<i>S. laxiflorum</i> ⁶	2.49	0.038	1220	K J	0.31	40	40	Heterosorghum
<i>S. leiocladum</i> ⁵	4.60	0.052	2254	F	2.30	20	10	Parasorghum
<i>S. macrospermum</i> ⁶	2.07	0.017	1014	L	0.26	40	40	Chaetosorghum
<i>S. matrankense</i> ⁶	2.51	0.029	1245	K J	1.26	10	10	Parasorghum
<i>S. nitidum</i> ⁵	8.79	0.060	4307	B	2.20	10, 20	20	Parasorghum
<i>S. plumosum</i> ⁵ (302489)	7.65	0.090	3748	C	1.28	10, 20, 30	30	Stiposorghum
<i>S. plumosum</i> ⁷ (302635)	10.30	0.249	5047	A	1.29	10, 20, 30	40	Stiposorghum
<i>S. propinquum</i> ⁵	1.52	0.012	745	MN	0.38	20	20	Eusorghum
<i>S. purpureosericeum</i>	4.18	0.022	2048	G	2.09	10	10	Parasorghum
<i>S. stipoides</i> ⁶	2.19	0.061	1073	L	1.10	10	10	Stiposorghum
<i>S. timorensis</i> ⁷	1.27	0.008	622	N	0.64	10, 20	10	Stiposorghum
<i>S. versicolor</i> (2x) ⁷	3.25	0.023	1592	I	1.62	10 ⁸	10	Parasorghum
<i>S. versicolor</i> (4x) ^{5,9}	6.67	0.086	3268	E	1.67	20	20	Parasorghum

¹ 1 pg = 980 mega base pairs (Mbp) (Cavalier-Smith, 1985).

² Duncan grouping for 2C DNA content (pg). Means followed by the same letter are not significantly different ($\alpha = 0.05$).

³ The genome size based on $x = 5$ was obtained using the following formula: genome size = [2C DNA content (pg)]/($2n$ chromosome no./5).

⁴ Reported chromosome numbers are from Lazarides *et al.* (1991) and Doggett (1988) unless referenced otherwise.

⁵ Calibration standard was *Lupinus texensis*; 2C DNA content is 2.44 pg.

⁶ Calibration standard was *Arabidopsis thaliana* ecotype Columbia; 2C DNA content = 314 Mbp or 0.32 pg (Bennett *et al.*, 2002).

⁷ Calibration standard was *Sorghum bicolor* TX623; 2C DNA content = 1.67 pg.

⁸ Chromosome number reported by Yu and Liang (1992).

⁹ Colchicine-induced autotetraploid (Sun *et al.*, 1994).

size. Chromosome numbers, with few exceptions, confirmed published counts (summarized in Lazarides *et al.*, 1991). A new chromosome number ($2n = 30$) was obtained for *S. amplus* (taxonomic identity confirmed by M. Lazarides) which was previously reported to be $2n = 10$. *Sorghum leiocladum* was reported to have $2n = 20$ chromosomes but a new number of $2n = 10$ was counted. The identity of the authors' accession of *S. leiocladum* was verified by senior taxonomist Ian Cowie at the Darwin Herbarium. Seven accessions of *S. nitidum* were counted, and all had $2n = 20$ chromosomes. Both $2n = 10$ and $2n = 20$ types of *S. nitidum* have been reported, but Garber (1950) questioned the reliability of the $2n = 10$ count. Multiple ploidy levels have been reported for *S. plumosum* ($2n = 10, 20, 30$) and *S. timorensis* ($2n = 10, 20$). The authors counted $2n = 30$ and 40 (new count) for *S. plumosum* and $2n = 10$ for *S. timorensis*.

DNA content

The DNA contents of the 21 *Sorghum* species are presented as picograms and as megabase pairs (Table 2). The 2C DNA content varies 8.1-fold among *Sorghum*

species from 1.27 to 10.30 pg (Table 2). Among the $2n = 10$ species, the 2C DNA content range is 3.6-fold (1.27–4.60 pg). The 2C DNA content among the $2n = 20$ species varies over a 5.8-fold range (1.52–8.79 pg). If the $2n = 20, 30$ and 40 species are polyploids, based on genomes of five chromosomes, then the DNA content may be analysed in terms of DNA content per $x = 5$ genome (Table 2). When this is done, DNA content per $x = 5$ genome encompasses an 8.8-fold range (0.26–2.30 pg).

Mean DNA content of annuals and perennials was compared by a *t*-test. For all *Sorghum* species, the mean DNA content of perennials ($2C = 6.20$ pg) is significantly greater ($\alpha = 0.01$) than the mean of the annuals ($2C = 2.92$ pg). However, when the same analysis was performed using adjusted $x = 5$ genome size, there was no significant difference in the means of the annuals (1.15 pg) and perennials (1.29 pg).

Mean DNA contents of native Australian species were compared by a *t*-test. For Australian species, the mean DNA content of perennials ($2C = 7.73$ pg) is significantly greater ($\alpha = 0.01$) than the mean of the annuals ($2C = 2.89$ pg). The same analysis performed on the adjusted $x = 5$ genome

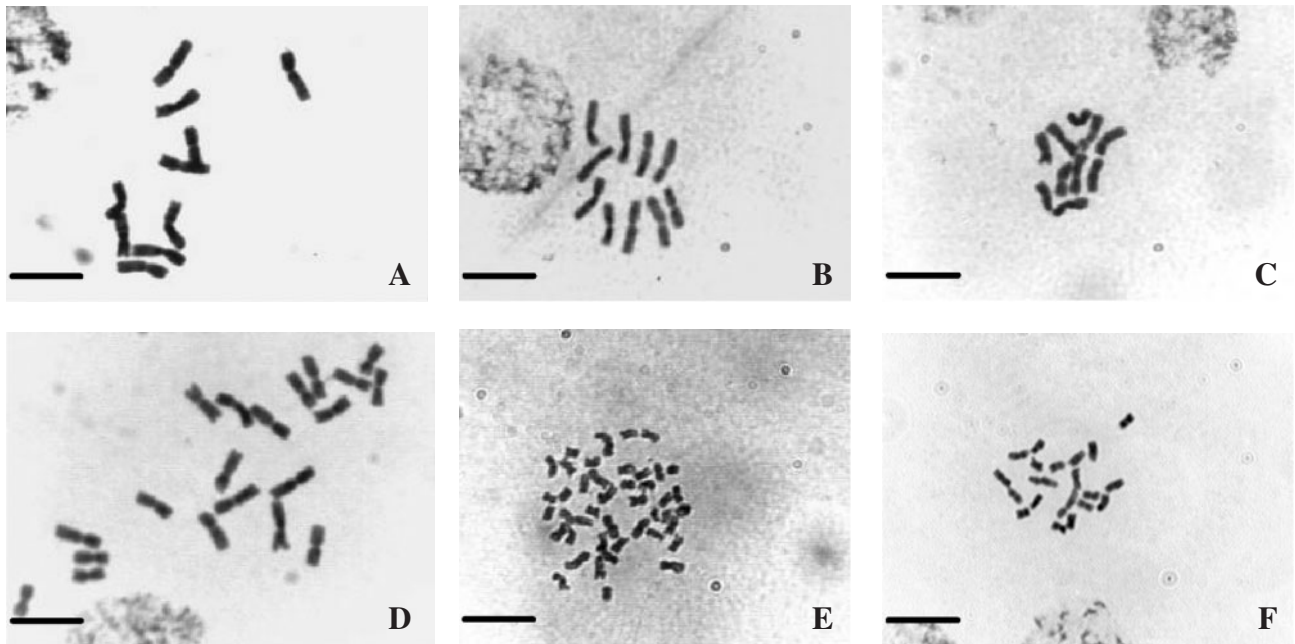


FIG. 1. Chromosomes of *Sorghum* species representing differences in number and size: (A) *S. leiocladum* ($2n = 10$); (B) *S. brachypodum* ($2n = 10$); (C) *S. timorense* ($2n = 10$); (D) *S. nitidum* ($2n = 20$); (E) *S. laxiflorum* ($2n = 40$); (F) *S. bicolor* ($2n = 20$). Scale bars = 10 μm .

size indicated a significant difference in the means of the annuals (1.09 pg) and perennials (1.66 pg) at $\alpha = 0.05$.

Figure 2 is a plot of 2C DNA content and the maximum southern latitude inhabited by each Australian species. Although there is a significant difference in mean maximum southern latitude (t -test; $\alpha = 0.01$) of the annual (-17.6 degrees) and perennial (-25.4 degrees) species, there is no significant correlation overall between maximum southern latitude and 2C DNA amount ($r = 0.307$; $P = 0.023$) or between maximum southern latitude and $x = 5$ DNA amount ($r = 0.407$; $P = 0.1015$).

Phylogenetic analysis

Figure 3 shows a strict consensus tree of six equally parsimonious solutions of 202 steps (CI = 0.792) for the 21 *Sorghum* species studied, based upon combined ITS/*ndhF* sequences. The 2C DNA content, $2n$ chromosome number and $x = 5$ genome size are indicated next to each species. The tree was rooted using *Zea mays*. There are two lineages for *Sorghum* (Fig. 3), one consisting of $2n = 10$ species and their polyploid relatives with large chromosomes (Fig. 1) and another containing $2n = 20$ and $2n = 40$ species with relatively small chromosomes (Fig. 1).

DISCUSSION

Chromosome number

The base chromosome number for the Andropogoneae has been considered to be either five or ten (Garber, 1950; Celarier, 1956). Garber (1950) stated that the base chromosome number could be $n = 10$ based upon the predominance of genera with a chromosome number of $n = 10$. Garber

(1950) also noted that the presence of genera with $n = 5$ suggests that five may be basic for the tribe, and questioned whether five represents the end point of a descending series in base chromosome numbers or the starting point of a polyploid series. Spangler *et al.* (1999) tried to resolve the base chromosome number for the Andropogoneae by superimposing chromosome numbers onto their phylogenetic tree based upon *ndhF* sequence analysis. The wide variety of taxa near the base of the tree with $n = 10$ led them to suggest that ten, not five, is the base number of the tribe.

The occurrence of both of $n = 5$ and $n = 10$ species of *Sorghum* raises questions regarding the base number of this genus. For example, there is genetic and molecular evidence that *Sorghum bicolor* ($n = 10$), although generally considered to be a diploid, may actually be a tetraploid. Brown (1943), Kidd (1952) and Endrizzi and Morgan (1955) reported bivalents in meiosis from haploid *S. bicolor*, indicating homology among chromosomes. Endrizzi and Morgan (1955) also observed translocations among progeny of haploids and proposed that these originated from recombination between homologous duplicated regions (resulting from polyploidy) in the haploid genome.

Several RFLP maps have been constructed for sorghum (Whitkus *et al.*, 1992; Berhan *et al.*, 1993; Chittenden *et al.*, 1994; Pereira *et al.*, 1994; Xu *et al.*, 1994; Dufour *et al.*, 1996; Peng *et al.*, 1999), and these have provided the basis for comparing the sorghum genetic map with that of other grass species. Comparison of the RFLP maps of sorghum and maize using maize-derived probes showed that many linkage groups are conserved between these two genomes (Whitkus *et al.*, 1992). Thirty-eight per cent of a common set of 89 RFLP probes for maize and sorghum were duplicated in sorghum and 72 % were duplicated in maize. Whitkus *et al.* (1992) suggested that the primary

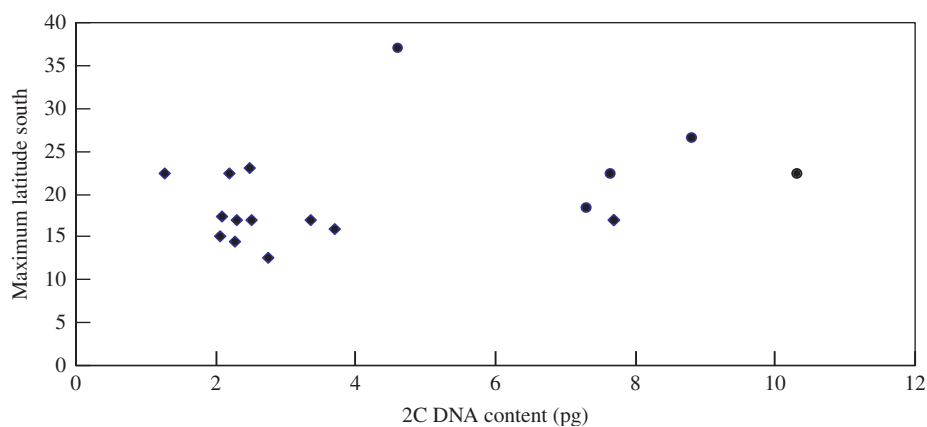


FIG. 2. 2C DNA content and the maximum southern latitude inhabited by native Australian *Sorghum* species. Diamonds represent annual and circles perennial species. The y-axis is the maximum latitude south of native Australian species.

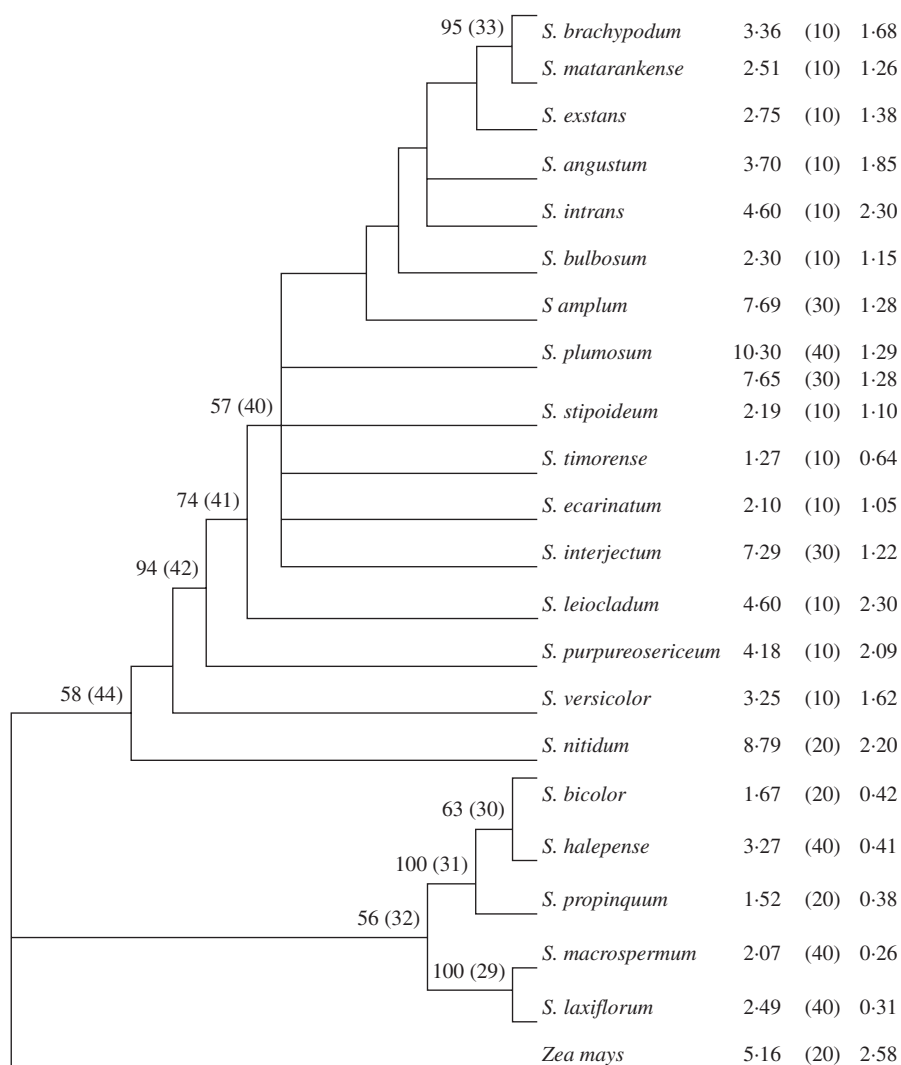


FIG. 3. The strict consensus tree of six equally parsimonious solutions of 202 steps (CI = 0.792). Bootstrap support (%) for various nodes from 10 000 replications is indicated above the corresponding node. Only bootstrap values of ≥ 50 are shown. Numbers in parentheses above each node represent unambiguous nucleotide substitutions. The tree shows bootstrap support, 56 % and 58 % respectively, for lineages consisting of (a) *S. bicolor* through *S. laxiflorum* and (b) *S. brachypodum* through *S. nitidum*. The tree was rooted using *Zea mays*. The 2C DNA content, 2n chromosome number (in parenthesis), and $x = 5$ genome size are denoted next to each species. The DNA content for corn (4C = 10.31 pg) is for inbred line Va35 (Laurie and Bennett, 1985).

processes involved in the divergence of maize and sorghum genomes were duplications (polyploidy or segmental), inversions, and intrachromosomal translocations. It was further suggested that, if maize and sorghum are both ancient polyploids, the larger proportion of duplicated loci in maize compared with sorghum might be due to a more rapid loss of duplicated segments in the evolution of sorghum (Whitkus *et al.*, 1992; Pereira *et al.*, 1994). Xu *et al.* (1994), Chittenden *et al.* (1994) and Peng *et al.* (1999) reported that between 8.2 % and 18.6 % of sorghum clones hybridized to duplicated loci. Peng *et al.* (1999) concluded that the distribution of duplicated loci in sorghum does not support the hypothesis that *S. bicolor* is of tetraploid origin.

Molecular cytogenetic evidence strongly supports a polyploid origin of sorghum. Gomez *et al.* (1998) and Zwick *et al.* (2000) detected by fluorescent *in situ* hybridization (FISH) a <280-bp tandemly repeated DNA sequence, CEN38, that differentially resided around the centromere of ten of the 20 chromosomes of *S. bicolor*. Gomez *et al.* (1998) proposed that the ten chromosomes displaying strong FISH signals composed one subgenome of a tetraploid, whereas the other ten chromosomes with little or no FISH signal represented another subgenome.

The sequence-based phylogenetic trees of Spangler *et al.* (1999) and that are presented in Fig. 3 leave unresolved the base chromosome number of the genus *Sorghum*. Spangler *et al.* (1999) showed chromosome numbers in an *ndhF* tree of 35 Andropogoneae species that included 12 species of *Sorghum*. There were two inaccuracies regarding chromosome numbers of *Sorghum* species in their report. First, they used a chromosome number of $n = 10$ for *S. leiocladum*. Here a chromosome number of $n = 5$ is reported for this species. Secondly, they included an unidentified species, *Sorghum* sp., in their analysis and assumed without documentation that its chromosome number was $n = 10$. When this species is removed and a correct chromosome number for *S. leiocladum* is used, the Spangler *et al.* (1999) tree has *Sorghum* split into two branches diverging from a node with equivocal chromosome numbers. One branch contains *Sorghum* with chromosome numbers of $2n = 20, 40$. The other branch contains the $2n = 10$ chromosome species.

The sequence phylogeny presented here, based upon combined ITS/*ndhF* sequences (Fig. 3), also has *Sorghum* split into two lineages, one with $2n = 10$ species with relatively large chromosomes and their related polyploids ($2n = 20, 30$ and 40) and a second containing $2n = 20$ and $2n = 40$ species with smaller chromosomes. The ancestral chromosome number of the genus *Sorghum* ($x = 5$ vs. $x = 10$) remains unresolved.

Nuclear DNA content

Nuclear DNA content is apparently important to the evolution and adaptation of plant species (Price, 1976, 1988; Bennett, 1973). DNA content affects cellular properties including nuclear volume, cell volume, the duration of mitosis and meiosis and minimum generation time (for reviews, see Price, 1976, 1988; Bennett, 1998), and may

influence ecological adaptation and distribution (Bennett, 1987; Bennett *et al.*, 2000). Geographic and ecological parameters including latitude, moisture availability and temperature, and growth form have been correlated with nuclear DNA content (Stebbins, 1966; Price, 1988; Bennett *et al.*, 2000). Genome size variation is common among congeneric species (Price, 1976), as is exemplified by the 3.6- and 5.8-fold range in 2C DNA content observed among the respective $2n = 10$ and $2n = 20$ *Sorghum* species reported herein.

Annual and perennial species of a genus often differ in DNA content (Price, 1976). For herbaceous angiosperms, the mean DNA content of perennials is greater than that of annuals (Bennett, 1972). For Australian native *Sorghum* species, the perennials have a significantly higher mean DNA content than annuals when comparing DNA content per 2C nucleus and for $x = 5$ genome size. Although there is geographical overlap in their distributions, the annual *Sorghum* species in Australia tend to occupy the lower (more tropical) latitudes with the higher subtropical to temperate latitudes (further south) inhabited by the perennials. A statistical analysis of DNA content and the maximum southern latitude at which the species naturally grow quantified this relationship. There is a significant difference ($\alpha = 0.01$) between the mean maximum latitude inhabited by the annual (-17.6 degrees) and perennial (-25.4 degrees) species (Fig. 2).

A current unresolved question concerns the relative frequency of increases and decreases in DNA content in angiosperm phylogeny (Bennetzen and Kellogg, 1997; Leitch *et al.*, 1998; Wendel *et al.*, 2002). Bennetzen and Kellogg (1997) analysed DNA content from a phylogenetic perspective in grasses and proposed that the evolution of DNA content in plants is primarily from low to high due to the combined effects of retroelement accumulation and polyploidy. They proposed that plants may have a 'one-way ticket to genomic obesity'. In contrast to an 'increase only' hypothesis for genome size evolution, it has been proposed that reduction in DNA content in plants has commonly occurred (Price, 1976, 1988; see also Bennetzen *et al.*, 2005). However, until recently this theory was lacking support based on well-founded phylogenetic relationships.

Wendel *et al.* (2002) analysed genome size evolution among species of the cotton tribe Gossypieae using a phylogenetic approach. They superimposed genome sizes on a well-supported sequence phylogeny and statistically inferred ancestral DNA contents. From this analysis it was concluded that both increases and decreases in DNA content have occurred repeatedly during evolution of the Gossypieae and DNA decreases actually exceeded increases. This supported the theory of a bi-directional dynamic nature of plant genome size evolution. Soltis *et al.* (2003) interpreted DNA content evolution in angiosperm phylogeny to be generally from low to high, but stated that genome size is dynamic with both increases and decreases occurring.

Superimposing genome sizes onto the combined ITS/*ndhF* sequence phylogeny (Fig. 3) provides a limited interpretation of the direction(s) of evolution of genome size in *Sorghum*. The lineage in Fig. 3 containing

S. brachypodium through *S. nitidum* is rooted with large $x = 5$ genome sizes, i.e. 2.20 pg (*S. nitidum*), 1.62 pg (*S. versicolor*), 2.09 pg (*S. purpureosericeum*) and 2.30 pg (*S. leiocladum*). This branch splitting *S. leiocladum* and the remaining species has moderate bootstrap support (74 %). The $x = 5$ genome sizes, with the exception of *S. brachypodium* (1.68 pg), *S. intrans* (2.3 pg) and *S. angustum* (1.85 pg), are all relatively low (0.63–1.38 pg). These species with smaller genomes probably represent evolutionary reductions in genome size, the extreme reduction having occurred in *S. timorense* (0.63 pg). Therefore, genome size evolution in the genus *Sorghum* apparently did not involve a 'one way ticket to genomic obesity' as was proposed for the grasses by Bennetzen and Kellogg (1997).

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